ld 3806-34-6 Date 15.12.2003

201-14967B

IUCLID

Data Set

Existing Chemical

CAS No.

: ID: 3806-34-6 : 3806-34-6

EINECS Name

: O,O'-dioctadecylpentaerythritol bis(phosphite)

EC No.

: 223-276-6

Molecular Formula

: C41H82O6P2

Status

Memo

: US HPV WESTON 618 Crompton Corp.

Printing date

: 15.12.2003

Revision date

Date of last update

: 15.12.2003

Number of pages

: 1

Chapter (profile)

: Chapter: 1, 2, 3, 4, 5, 6, 7, 8, 10

Reliability (profile)

: Reliability: without reliability, 1, 2, 3, 4

Flags (profile)

: Flags: without flag, confidential, non confidential, WGK (DE), TA-Luft (DE), Material Safety Dataset, Risk Assessment, Directive 67/548/EEC, SIDS

2. Physico-Chemical Data

ld 3806-34-6 **Date** 15.12.2003

2.1 MELTING POINT

Value

37 - 46 °C

Sublimation

Method

•

Year

GLP

: no data

Test substance

Chemical name: Distearyl Pentaerythritol Diphosphite

CAS No.: 3806-34-6

Trade name: Weston 618F, 618G Phospites Purity: No data, likely to be technical grade

Reliability

: (4) not assignable

Manufacturer's technical data sheet

22.10.2003

(2)

2.2 BOILING POINT

Value

705 °C at

Decomposition

:

Method

other: Calculated using MPBPWIN v 1.40

Year

2003

GLP

:

Test substance

Chemical name: Distearyl Pentaerythritol Diphosphite

CAS No.: 3806-34-6

Reliability

: (2) valid with restrictions

17.11.2003

(7)

2.4 VAPOUR PRESSURE

Value

1.06E-18 hPa at 25 °C

Decomposition

.

Method

other (calculated): MPBPWIN v 1.40

Year

2003

GLP

. 200

Test substance

: Chemical name: Distearyl Pentaerythritol Diphosphite

CAS No.: 3806-34-6

Reliability

: (2) valid with restrictions

18.11.2003

(7)

2.5 PARTITION COEFFICIENT

Partition coefficient

octanol-water

Log pow

15 at °C

pH value Method

other (calculated): KOWWIN v 1.66

Year

offici (calculated). NOVVVVI

GLP

: 2003

Test substance

: Chemical name: Distearyl Pentaerythritol Diphosphite

CAS No.: 3806-34-6

Reliability 22.10.2003

: (2) valid with restrictions

2. Physico-Chemical Data

Id 3806-34-6Date 15.12.2003

2.6.1 SOLUBILITY IN DIFFERENT MEDIA

Solubility in : Water Value : at °C

pH value

concentration : at °C

Temperature effects

Examine different pol.

pKa : at 25 °C

Description

Stable

Deg. product

Method : other: Calculated using WSKOW v 1.40

Year : 2003

GLP

Test substance: Chemical name: Distearyl Pentaerythritol Diphosphite

CAS No.: 3806-34-6

Result : Water solubility = 2.95E-12 mg/L

Reliability : (2) valid with restrictions

18.11.2003

3. Environmental Fate and Pathways

ld 3806-34-6 **Date** 15.12.2003

3.1.1 PHOTODEGRADATION

Type : air Light source :

Light spectrum: nm

Relative intensity : based on intensity of sunlight

INDIRECT PHOTOLYSIS

Sensitizer : OH

Conc. of sensitizer : 1500000 molecule/cm³

Rate constant : .000000001863 cm³/(molecule*sec)

Degradation: % after

Deg. product

Method : other (calculated): AOP v 1.90

Year : 2003

GLP :

Test substance : Chemical name: Distearyl Pentaerythritol Diphosphite

CAS No.: 3806-34-6

Result : T1/2 = 0.689 hours

22.10.2003 (7)

3.1.2 STABILITY IN WATER

3.3.1 TRANSPORT BETWEEN ENVIRONMENTAL COMPARTMENTS

Type : fugacity model level III

Media

Air : % (Fugacity Model Level I)
Water : % (Fugacity Model Level I)
Soil : % (Fugacity Model Level I)
Biota : % (Fugacity Model Level II/III)
Soil : % (Fugacity Model Level II/III)

Method : other: Calculation using EPIWIN Level III Fugacity Model

Year : 2003

Test condition: Henry's Law Constant: 8.15E-6 atm-m3/mole (Henrywin program)

Vapor pressure: 8E-17 mmHg (Mpbpwin program)

MPt.: 46°C (user entered)

Log Kow: 15.1 (Kowwin program) Soil Koc: 4.6E+14 (calc by model)

1000 kg/hr emissions to air, water and soil compartments.

Test substance : Chemical name: Distearyl Pentaerythritol Diphosphite

CAS No.: 3806-34-6

	Mass Amount	Half-life	Emissions
	(percent)	(hr)	(kg/hr)
Air	0.02	1.38	1000
Water	2.39	1440	1000
Soil	28.6	1440	1000
Sediment	68.9	5760	0

	Fugacity (atm)	Reaction (kg/hr)	Advection (kg/hr)	Reaction (percent)	Advection (percent)
Air	8.51E-20	926	18.4	30.9	0.614

3. Environmental Fate and Pathways

ld 3806-34-6 Date 15.12.2003

Soil Sediment 1.22E-22 1.32E-20 1050 631

0 105 35 21

0 3.5

Persistence time: 2540 hr Reaction time: 2820 hr Advection time: 24900 hr Percent reacted: 89.8 Percent advected: 10.2

Half-lives (hr), (based upon Biowin (ultimate) and Aopwin):

Air: 1.378 Water: 1440 Soil: 1440 Sediment: 5760

Biowin estimate: 1.964 (months)

Advection times (hr):

Air: 100 Water: 1000 Sediment: 5E+4

Reliability

: (1) valid without restriction

22.10.2003

(7)

BIODEGRADATION

Type

aerobic

Inoculum

Deg. product

Method

: other: calculated using Biowin v 4.0

Year

2003

GLP

Test substance

Result

MITI Linear Biodegradation Probability = 0.3588 MITI Non-linear Biodegradation Probability = 0.0660

The substance is predicted to be not readily biodegradable

Test substance

Chemical name: Distearyl Pentaerythritol Diphosphite

CAS No.: 3806-34-6

Reliability

(2) valid with restrictions

22.10.2003

4. Ecotoxicity

ld 3806-34-6

Date 15.12.2003

ACUTE/PROLONGED TOXICITY TO FISH 4.1

Type

Species Exposure period

96 hour(s)

Unit

mg/l

Method

other: Calculated using ECOSAR v 0.99g

Year

2003

GLP

Test substance

Chemical name: Distearyl Pentaerythritol Diphosphite

CAS No.: 3806-34-6

Result

: LC50 = 2.94E-10 mg/L

The LC50 value is above the estimated water solubility of this substance.

Reliability

: (2) valid with restrictions

22.10.2003

(7)

4.2 **ACUTE TOXICITY TO AQUATIC INVERTEBRATES**

Type

Species

Daphnia sp. (Crustacea)

Exposure period

48 hour(s)

Unit

mg/l other: Calculated using ECOSAR v 0.99g

Method Year

2003

GLP

Test substance

Chemical name: Distearyl Pentaerythritol Diphosphite

CAS No.: 3806-34-6

Result

: LC50 = 7.76E-10 mg/L

The LC50 value is above the estimated water solubility of this substance.

Reliability

(2) valid with restrictions

22,10,2003

(7)

4,3 TOXICITY TO AQUATIC PLANTS E.G. ALGAE

Species

Endpoint

Exposure period

Test substance

96 hour(s) mg/l

Unit Method

other: Calculated using ECOSAR v 0.99g

Year

2003

GLP

Chemical name: Distearyl Pentaerythritol Diphosphite

CAS No.: 3806-34-6

Result

: EC50 = 1.03E-09 mg/L

The EC50 value is above the estimated water solubility of this substance.

Reliability 22.10.2003 (2) valid with restrictions

5. Toxicity Id 3806-34-6
Date 15.12.2003

5.1.1 ACUTE ORAL TOXICITY

Type : LD50

Value : > 10000 mg/kg bw

Species : rat

Strain : Sherman Sex : male/female

Number of animals : 10

Vehicle : other: vegetable oil Doses : 10,000 mg/kg

Method : other: US Testing Co., Inc. Method

Year : 1971 **GLP** : no

Test substance : Chemical name: Distearyl Pentaerythritol Diphosphite

CAS No.: 3806-34-6 Trade name: Weston 618

Purity: No data, likely to be technical grade

Result : No. of deaths: 0

Clinical signs: None of the animals showed any signs of toxicity at the

maximum dose that could be given at a single administration

Test condition : Weight of animals: 200 - 220 g

Concentration administered: Test material suspended in vegetable oil at a

ratio of 2:5 grams of sample/mL of oil Administration method: Gavage Post dose observation period: 14 days

Reliability : (2) valid with restrictions

Apparently well-conducted study

20.10.2003 (6)

5.1.3 ACUTE DERMAL TOXICITY

Type : LD50

Value : > 2000 mg/kg bw

Species : rabbit

Strain : New Zealand white

Sex : male/female

Number of animals : 10

Vehicle :

Doses : 2000 mg/kg

Method : OECD Guide-line 402 "Acute Dermal Toxicity"

 Year
 : 1994

 GLP
 : yes

Test substance : Chemical name: Distearyl Pentaerythritol Diphosphite

CAS No.: 3806-34-6

Trade Name: Weston W618F

Lot No.: HBA242

Purity: No data, likely to be technical grade

Result : Mortality: No deaths during the study

Clinical observations: 2 females had soft stool on day 1. Two rabbits had their colars caught in their mouth during test material exposure and one of these animals had wet red material around the mouth. There were no

Id 3806-34-6

Date 15.12.2003

Dermal observations: The test material induced very slight to moderate erythema on all rabbits and very slight edema on eight rabbits.

Desquamation was present on 8 sites by day 7 and one site by day 14. There were no other dermal findings. Three sites had very slight erythema and/or desquamation at study termination (day 14).

Body weights: No remarkable changes or differences in body weights noted during this study.

Necropsy: Accessory splenic tissue, a common congenital abnormality in this strain of rabbit was noted for 5 animals. There were no other gross necropsy findings for all examined tissues.

Test condition : Age: Approximately 11 weeks old

Weight: 2098 - 2240 g

Volume administered or concentration: Applied neat

Post dose observation period: 14 days

Reliability : (1) valid without restriction

Guideline study conducted to GLP

20.10.2003 (8)

5.5.0. ?@?@ • EYE IRRITATION

Species : rabbit Concentration : 10 %

Dose : other: unspecified Exposure time : unspecified

Comment

Number of animals : 6

Vehicle : other: Cottonseed oil Result : slightly irritating

Classification : not irritating

Method : other: Federal Register, Vol 29, No. 182, p 13009, 17 September 1964

Year : 1971 **GLP** : no

Test substance : Chemical name: Distearyl Pentaerythritol Diphosphite

CAS No.: 3806-34-6

Trade name: Weston 618 Phosphite Purity: No data, likely to be technical grade

Result : The test material produced a very mild conjunctival effect in two of the

animals which cleared by the second day of observation

22.10.2003 (1)

5.5 REPEATED DOSE TOXICITY

Туре

Species : rat

Sex : male/female

Strain : other: Charles River albino

Route of admin. : oral feed Exposure period : 90 days Frequency of treatm. : daily ad libitum

Post exposure period : none

Doses : 300, 1,000, 3,000 ppm

5. Toxicity Id 3806-34-6 Date 15.12.2003

NOAEL

: > 3000 ppm

Method

: other: Industrial Bio-Test Laboratories Inc. test method

Year GLP 1972 no

Test substance

Chemical name: Distearyl Pentaerythritol Diphosphite

CAS No.: 3806-34-6

Trade name: Weston Phosphite 618

Lot no.: 24

Purity: No data, likely to be technical grade

Result

Body weight: Statistical comparisons of final body weights and total weight gains revealed no significant differences between test and control rats

Food/water consumption: Test rats ate amounts of food comparable to that consumed by control rats.

Clinical signs: No untoward behavioral reactions were noted among any of the animals employed in the study.

Ophthalmologic findings: Non reported

Hematologic findings: No outstanding differences between test and control rats were noted with respect to any of the parameters investigated (hematocrit value, erythrocyte count, hemoglobin concentration, total leukocyte count, differential leukocyte count).

Clinical blood chemistry findings: Values for blood urea nitrogen concentration, serum alkaline phosphatase activity, serum glutamic-pyruvic transaminase activity and fasted blood glucose concentration for test rats compared well with controls.

Urine analysis: No significant differences between the urine of test rats and control rats were observed when urine was analysed for glucose concentration, albumin concentration, pH, specific gravity and microscopic elements examination.

Mortality and time to death: Six deaths occurred during the study. All of these deaths resulted from trauma incurred during the collection of blood samples. These deaths occurred in the control as well as the test groups and were not attributed to the ingestion of the test material.

Gross pathology: No outstanding differences were noted between test and control rats.

Organ weight changes: The only statistically significan difference reported was the liver/body weight ratio for the 3000 ppm males. The authors of the study concluded that the lack of any consistent dietary or sex-related response indicates that the intergroup differences were not related to treatment.

Histopathology: All of the lesions noted in the microscopic examination of tissues were those of spontaneous disease and are not unusual for the albino rat. The most frequent findings were lesions in the trachea and lungs, indicating chronic murine pneumonia. These occurred in the control as well as the treated rats.

Test condition

Test subjects

Age at study initiation: no data

Mean body weight at study inititiation: 99 g (male), 115 g (female)

ld 3806-34-6

Date 15.12.2003

Study Design

Vehicle: Standard rat ration

Clinical observations performed and frequency:

Body weight: Measured on the first day of the test and at weekly intervals thereafter. Analysed statistically at the end of the study.

Food consumption: Data were collected individually for five rats of each sex in every group weekly during the study.

Abnormal reactions and death: Recorded daily during the investigation.

Blood and urine: Samples were collected individually from 10 rats of each sex from both the control and the 3,000 ppm groups after 45 and 84 days of feeding for analysis.

Organs examined at necropsy (macroscopic and microscopic): Esophagus, stomach (cardia, fundus and pylorus), small intestine (duodenum, jejunum and ileum), cecum, colon, liver, kidneys, spleen, pancreas, urinary bladder, pituitary gland, adrenal gland, testes, seminal vesicle, ovary, bone marrow, thyroid gland, parathyroid gland, salivary gland, prostate gland, heart, aorta, lung, lymph node (cervical and mesenteric), skeletal muscle, peripheral nerve, bone (femur), spinal cord, uterus, trachea, eye, optic nerve and brain (cerebrum, cerebellum and pons)

Organ weights: Statistical analyses were conducted upon the absolute organ weights and their corresponding ratios to the weight of the body and brain. An Analysis of Variance was conducted first and any significant effects disclosed by that treatment were further studied by t-tests.

Reliability

: (1) valid without restriction

Well conducted and reported study

20.10.2003

(4)

5.5 GENETIC TOXICITY 'IN VITRO'

Type

Ames test

System of testing

: Salmonella typhimurium strains TA97, TA98, TA100, TA102

Escherichia coli strain WP2/pKM101

Test concentration

0, 0.05, 0.1, 0.2, 9.5 mg/plate

Cycotoxic concentr. Metabolic activation

with and without

Result

negative

Method

other: Maron & Ames (1983)

Year **GLP**

1985

no data

Test substance

Chemical name: Distearyl Pentaerythritol Diphosphite

CAS No.: 3806-34-6

Result

Dispersing the test substance into solutions of acetone/Tween-80 at doses of 0.2mg - 0.5 mg/plate produced cloudy solutions.

Under these conditions, there was neither an increase in the number of revertant cells, nor any toxicity.

It was judged that the test was negative under these conditions.

Reliability

(4) valid without restriction Summary of study only available

5. **Toxicity**

ld 3806-34-6 Date 15.12.2003

15.12.2003

(5)

GENETIC TOXICITY 'IN VIVO'

Type

Micronucleus assay

Species Sex

mouse male/female

Strain

: ICR

Route of admin.

: i.p.

Exposure period

: 24, 48 hours

Doses

500, 1000, 2000 mg/kg

Result

: negative

Method

OECD Guide-line 474 "Genetic Toxicology: Micronucleus Test"

Year

2003

GLP

yes

Test substance

Chemical name: Distearyl Pentaerythritol Diphosphite CAS No.: 3806-34-6

Trade name: Weston 618F

Lot No.: H41425

Result

Effect on mitotic index or PCE/NCE ratio by dose level by sex: See table

below.

Genotoxic effects: Negative

Mortality at each dose level by sex:

Pilot toxicity study: No mortality occurred at any dose, up to the maximum tested of 2000 mg/kg.

Main study: No mortality occurred at any dose level during the course of the study.

Clinical signs:

Pilot toxicity study: Piloerection was seen in male mice at 100 and 1000 mg/kg and in male and female mice at 2000 mg/kg and lethargy in males at 1000 mg/kg and in male and female mice at 2000 mg/kg.

Main study: Lethargy was observed in male and female mice at 1000 and 2000 mg/kg and piloerection in males and females at all doses tested. All other mice treated with test or control articles appeared normal during the course of the study.

Bodyweight changes:

Pilot toxicity study: Change in group mean bodyweights ranged from -2.9% (male, 2000 mg/kg) to +0.4% (female, 2000 mg/kg) after 3 days.

Mutant/aberration/mPCE/polyploidy frequency, as appropriate: See table below

Food/water consumption: no data available

Table: Summary of Bone Marrow Micronucleus analysis

Treatment	Sex	Time	No. of	PCE/Total	Change from	Micronucleated Polychromatic Erythrocytes	
(20mL/kg)		(hr)	mice	Erythrocytes	Control (%)	Number per 1000 PCEs	Number per
				(mean ± SD)		(mean ± SD)	PCEs Scored ¹

5. Toxicity

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	F	24	5	0.526±0.09	-	0.5±0.35	5/ 10000
Test article							
500 mg/kg	M	24	5	0.451±0.03	-1	0.6±0.22	6/ 10000
	F	24	5	0.465±0.02	-12	0.5±0.50	5/ 10000
1000 mg/kg	M	24	5	0.473±0.04	4	0.5±0.35	5/ 10000
	F	24	5	0.479±0.05	-9	0.5±0.35	5/ 10000
2000 mg/kg	M	24	5	0.447±0.03	-2	0.5±0.35	5/ 10000
	F	24	5	0.485±0.06	8	0.7±0.27	7/ 10000
$\mathbb{C}\mathbb{P}^2$	M	24	5	0.335±0.03	-27	22.2±2.20	*222/ 10000
50 mg/kg	F	24	5	0.325±0.01	-38	20.4±2.43	*204/ 10000
Corn oil	M	48	5	0.502±0.06	_	0.3±0.27	3/ 10000
	F	48	5	0.483±0.05	-	0.6±0.22	6/ 10000
Test article							
2000 mg/kg	M	48	5	0.471±0.05	-6	0.6±0.22	6/ 10000
	F	48	5	0.467±0.05	-3	0.8±0.27	8/ 10000

¹*statistically significant, p<=0.05 (Kastenbaum-Bowman Tables).

Test condition

Age at study initiation: 6 - 8 weeks old at the initiation of each phase of the study.

No. of animals per dose:

Pilot toxicity study: 2 male mice dosed at 1, 10, 100 or 1000 mg/kg b.w.; 5 male and 5 female mice dosed at 2000 mg/kg.

Main study: Groups of 5 male/5 female mice dosed at 0, 500, 1000, 2000 mg/kg (euthanized at 24 h); Groups of 5 male/5 female dosed at 0, 2000 mg/kg (euthanized at 48 h).

Route: i.p.

Vehicle: Corn oil.

Controls: Vehicle (Corn oil), cyclophosphamide monohydrate (positive).

Clinical observations performed: Clinical signs, mortality, bodyweight

Organs examined at necropsy: none

Criteria for evaluating results: The incidence of micronucleated polychromatic erythrocytes per 2000 polychromatic erythrocytes was determined for each mouse and treatment group. Statistical significance was determined using the Kastenbaum-Bowman tables which are based on the binomial distribution. In order to quantify the proliferation state of the bone marrow as an indicator of bone marrow toxicity, the proportion of polychromatic erythrocytes to total erythrocytes was determined for each animal and treatment group. The test article was considered to induce a positive response if a dose-responsive increase in micronucleated polychromatic erythrocytes was observed and one or more doses were statistically elevated relative to the vehicle control (p<=0.05, Kastenbaum-Bowman Tables) at any sampling time. However, values that were statistically significant but did not exceed the range of historical negative or vehicle controls were judged as not biologically significant. The test article was judged negative if no statistically significant increase in micronucleated polychromatic erythrocytes above the concurrent vehicle control values and no evidence of dose responses were observed at any sampling time.

Criteria for selection of M.T.D.: based on preliminary toxicity study.

(1) valid without restriction

Reliability

² cyclophosphamide monohydrate

5. Toxicity

ld 3806-34-6 **Date** 15.12.2003

5.8.1 TOXICITY TO FERTILITY

5.8.2 DEVELOPMENTAL TOXICITY/TERATOGENICITY

(1) Food and Drug Research Laboratories, Inc. (1971) Weston 618 Phosphite Rabbit Eye Irritation Study, Report No: IBL 10201-F GE Specialty Chemicals, Inc. (2000). Weston 618F, 618G Phosphites, Technical Data (2) Sheet CA-200H Gudi, R., & Krsmanovic, L. (2003) Bioreliance, Mammalian erythrocyte micronucleus test, (3) Study No. AA77XC.123.BTL Industrial Bio-Test Laboratories, Inc. (1972), 90-day subacute oral toxicity study with (4) Weston Phosphite 618 in albino rats, Report No. B1704. Takizawa, Y (1984) Public hygenic Section, Medical Dept., Akita University, Japan, Report (5) No. BWCT-022-5 United States Testing Company, Inc. (1971). Report of Test Number 51044. (6)US EPA, EPIWIN v3.10, EPI Suite Software, 2000 (7) Wil Research Laboratories, Inc. (1994), Acute Dermal Toxicity Study of Weston W618F in (8) Albino Rabbits, Report No. WIL-202008

ld 3806-34-6 **Date** 15.12.2003

9. References